1	Fussing about fission: defining variety among mainstream and
2	exotic apicomplexan cell division modes
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29 Keywords: Apicomplexa, cell division, cell cycle, karyokinesis, schizogony, endodyogeny,

- 30 endopolygeny, binary fission
- 31

32 Abstract

33 Cellular reproduction defines life, yet our textbook-level understanding of cell division is limited 34 to a small number of model organisms centered around humans. The horizon on cell division 35 variants is expanded here by advancing insights on the fascinating cell division modes found in 36 the Apicomplexa, a key group of protozoan parasites. The Apicomplexa display remarkable 37 variation in offspring number, whether karyokinesis follows each S/M-phase or not, and whether 38 daughter cells bud in the cytoplasm or bud from the cortex. We find that the terminology used to 39 describe the various manifestations of asexual apicomplexan cell division emphasizes either the 40 number of offspring or site of budding, which are not directly comparable features and has led to 41 confusion in the literature. Division modes have been primarily studied in two human pathogenic Apicomplexa, malaria-causing Plasmodium spp. and Toxoplasma gondii, a major cause of 42 43 opportunistic infections. *Plasmodium* spp. divide asexually by schizogony, producing multiple 44 daughters per division round through a cortical budding process, though at several life-cycle 45 nuclear amplifications are not followed by karyokinesis. T. gondii divides by endodyogeny 46 producing two internally budding daughters per division round. Here we add to this diversity in 47 replication mechanisms by considering the cattle parasite *Babesia bigemina* and the pig parasite 48 Cystoisospora suis. B. bigemina produces two daughters per division round by a 'binary fission' 49 mechanism whereas C. suis produces daughters through both endodyogeny and multiple internal 50 budding known as endopolygeny. In addition, we provide new data from the causative agent of 51 equine protozoal myeloencephalitis (EPM), Sarcocystis neurona, which also undergoes 52 endopolygeny but differs from C. suis by maintaining a single multiploid nucleus. Overall, we 53 operationally define two principally different division modes: internal budding found in cyst-54 forming Coccidia (comprising endodyogeny and two forms of endopolygeny) and external 55 budding found in the other parasites studied (comprising the two forms of schizogony, binary 56 fission and multiple fission). Progressive insights into the principles defining the molecular and 57 cellular requirements for internal versus external budding, as well as variations encountered in 58 sexual stages are discussed. The evolutionary pressures and mechanisms underlying 59 apicomplexan cell division diversification carries relevance across Eukaryota.

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60 **Contribution to the Field**

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62 Mechanisms of cell division vary dramatically across the Tree of Life, but the mechanistic basis has only been mapped for several model organisms. Here we present cell division strategies 63 64 across Apicomplexa, a group of obligate intracellular parasites with significant impact on 65 humans and domesticated animals. Asexual apicomplexan cell division is organized around 66 assembly of daughter buds, but division forms differ in the cellular site of budding, number of offspring per division round, whether each S-phase follows karvokinesis and if mitotic rounds 67 68 progress synchronously. This varies not just between parasites, but also between different life-69 cycle stages of a given species. We discuss the historical context of terminology describing 70 division modes, which has led to confusion on how different modes relate to each other. 71 Innovations in cell culture and genetics together with light microscopy advances have opened up 72 cell biological studies that can shed light on this puzzle. We present new data for three division 73 modes barely studied before. Together with existing data, we show how division modes are 74 organized along phylogenetic lines and differentiate along external and internal budding 75 mechanisms. We also discuss new insights into how the variations in division mode are regulated 76 at the molecular level, and possess unique cell biological requirements.

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77 1 Introduction

78 Reproduction is critical for perpetuating a species and lies at the core of the definition of life. 79 Yet, the modes by which cell division can occur are very diverse. The molecular and cellular 80 mechanisms underlying such differences have been dissected for a limited set of model 81 organisms, most of which carry resemblance to mammalian/human cell division. The 82 Apicomplexa comprise a protozoan phylum harboring human pathogens, like malaria-causing 83 Plasmodium spp., opportunistic parasites like Toxoplasma gondii and Cryptosporidium spp., and 84 emerging pathogens like Babesia spp. (phylogeny in Fig 1). In addition, many are of economic 85 relevance in agriculture and companion animals such as *Babesia* spp. *Theileria* spp. and *Eimeria* 86 spp. Furthermore, many Apicomplexa infect other birds, mammals, reptiles, amphibians, fish, 87 and invertebrates, but their cell biology has only been studied minimally. The asexual 88 multiplication cycles of the Apicomplexans are diverse, and their sexual cycles add even more 89 diversity. The well-studied Apicomplexa relevant to humans display a diverse spectrum of 90 asexual cell division strategies both within and among species, providing an excellent 91 opportunity for comparative biology. Indeed, we posit that by studying the principles in more 92 distant organisms such as protozoa, which span a wide breadth of evolutionary history and 93 correspondingly diverse biology, we can obtain fascinating new insights and principles. 94 Asexual apicomplexan cell division revolves around variations in budding through the 95 assembly of a membrane skeleton that ultimately underlies the plasma membrane. The beginning- and end-point of a cell division round across all Apicomplexa is a "zoite", a cell type 96 97 with the phylum-defining complex of apical secretory organelles and cytoskeletal structures 98 (Leander and Keeling, 2003) (e.g. Fig 2A, 3F1, 4A, 5E1, 6C1). These apical structures uniformly 99 facilitate host-cell invasion, an essential step in the obligate intracellular life style of 100 apicomplexan parasites (Gubbels and Duraisingh, 2012; Sharma and Chitnis, 2013; Frenal et al., 101 2017a). The cortical cytoskeleton just below the plasma membrane, however, plays an additional 102 role as a key structure facilitating budding across asexual division modes for Apicomplexa 103 (Anderson-White et al., 2012; Kono et al., 2016). From the outside in, it is built up from three 104 uniformly conserved elements: alveolar vesicles (alveoli) that anchor the myosin motor enabling 105 motility (Frenal et al., 2017a), an epiplastin (a.k.a. alveolar) protein meshwork (Goodenough et 106 al., 2018), and a series of sub-pellicular, longitudinal microtubules emanating from the apical 107 end. The number and length of the microtubules vary across parasite species and stage (Spreng et

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108 al., 2019), but the alveoli and epiplastin meshwork are universally conserved and make up the 109 inner membrane complex (IMC) (Kono et al., 2012). The epiplastin family proteins contain Val-110 Pro-Val (VPV) repeats and are generally known as alveolins or IMC proteins (Gould et al., 2008; 111 Anderson-White et al., 2011; Kono et al., 2012; Al-Khattaf et al., 2015). Cytoskeletal assembly 112 initiates from the centrosome and all three elements are simultaneously assembled in an apical to 113 basal direction (Chen and Gubbels, 2013; Francia and Striepen, 2014; Suvorova et al., 2015). 114 Another important and shared feature is that across asexual division modes, budding appears to 115 be tied always to a round of S-phase and mitosis (Francia and Striepen, 2014; Suvorova et al., 116 2015). A fourth cytoskeleton element that is conserved across division modes is the basal 117 complex, a ring residing at the basal end of the daughter bud (Ferguson et al., 2008). This ring 118 contains MORN1, a scaffolding protein that is essential for maintenance of the daughter bud's 119 basal end integrity (Gubbels et al., 2006; Hu, 2008; Heaslip et al., 2010; Lorestani et al., 2010; 120 Kono et al., 2016). Proteins with repetitive MORN domains are often found in cilia and flagella 121 (Mecklenburg, 2007; Shetty et al., 2007; Tokuhiro et al., 2008; Morriswood and Schmidt, 2015); 122 indeed all apicomplexan division modes share several features with cilia or flagella assembly as 123 seen across eukaryotes. For example, striated fiber assembly (SFA) fibers are found in flagellar 124 assembly in algae to orient the basal body in the flagellum, which in Apicomplexa anchor the 125 centrosome in the daughter bud (Francia et al., 2012). A spindle assembly abnormal protein 6 126 (SAS6)-like protein found in flagellar basal bodies is also found in the apical polar ring (APR). 127 the microtubule organizing center (MTOC) nucleating the subpellicular microtubules (and 128 conoid, if present) (de Leon et al., 2013; Francia et al., 2015; Wall et al., 2016). Finally, the base 129 of cilia and flagella is often decorated with a contractile centrin and a myosin (Roberts et al., 130 2004; Trojan et al., 2008), which, at least in T. gondii, are represented by TgCentrin2 and 131 Myosin I and J (Hu et al., 2006; Hu, 2008; Frenal et al., 2017b). 132 Despite this central plan, many variations exist on what precedes daughter budding, and how 133 daughter bud formation is orchestrated. The first variable is the number of offspring per mother 134 cell in an asexual apicomplexan division round, which can be as few as two or nearly 100,000. 135 The principle behind the uncoupling of daughter formation from S/M-phase was recently 136 established in T. gondii by the presence of a bipartite centrosome: the centrosome's inner-core 137 proximal to the nucleus regulates the S/M cycle independently of the centrosome's outer-core

distal from the nucleus, which organizes daughter bud formation (Chen and Gubbels, 2013;

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139 Suvorova et al., 2015; Chen and Gubbels, 2019). Moreover, cyclins and cyclin-dependent 140 kinases (CDKs), as found in higher eukaryotes, ultimately control cell cycle progression but their 141 nature and modes of operation are specifically tailored to the needs of different apicomplexan 142 cell cycle variation (e.g. (Roques et al., 2015; Alvarez and Suvorova, 2017; Ganter et al., 2017; 143 Naumov et al., 2017), recently reviewed in (Matthews et al., 2018; White and Suvorova, 2018)). 144 A second variable is whether karyokinesis follows each S-phase and mitosis (S/M-phase) or not. 145 Finally, daughters can either assemble within the mother's cytoplasm (internal budding), or 146 assemble and emerge directly from the plasma membrane of the mother cell (cortical or 147 peripheral budding). Strikingly, these modes can vary not only between Apicomplexa, but also 148 across different life-cycle stages within a single species. 149 Although these insights describe the key principles, the details on cell biological mechanisms 150 facilitating the various division modes are intermittent and often compounded by historical 151 terminology which either do not accurately capture the shared principles or clearly define 152 differences. Here we highlight the best-understood cell division modes, schizogony in

153 *Plasmodium* spp. and endodyogeny in *T. gondii*, supplemented with new data on emerging

systems displaying different forms of endopolygeny in *Cystoisopora suis* and *Sarcocystis*

neurona, as well as binary fission in *Babesia bigemina*. Emerging insights are subsequently used

- 156 to assess their kinship and to chart principles underlying these distinct cell division modes.
- 157

158 2 Material and Methods

159 2.1 Babesia spp. culture

160 B. bigemina strain JG-29 (Mexico) was kindly provided by Dr. David Allred (University of 161 Florida) and grown as described previously (Vega et al., 1985) with modifications as described 162 here. Parasites were culture adapted over a month to grow efficiently in tissue culture media 163 RPMI-1640 supplemented with 25 mM HEPES, 50 mg/l hypoxanthine, 2.42 mM sodium 164 bicarbonate, and 4.31 mg/ml AlbuMAX-II (Invitrogen). Before addition of AlbuMAX-II and 165 sodium bicarbonate, the pH was adjusted to 6.75. Parasites were grown under microaerophilous 166 stationary phase culture conditions with a hypoxic atmosphere of 1:5:94 oxygen: carbon dioxide: 167 nitrogen at 4% hematocrit in washed, defibrinated bovine red blood cells (Hemostat Labs, Dixon, 168 CA).

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169	B. divergens strain Rouen 1987, a kind gift from Drs. Kirk Deitsch and Laura Kirkman
170	(Weill Cornell Medical College), was grown in human erythrocytes as described (Paul et al.,
171	2016). Human erythrocytes were purchased from Research Blood Components (Boston, USA).
172	
173	2.2 <i>Cystoisospora suis</i> culture
174	Oocysts of C. suis strain Wien-I (Austria) were isolated from porcine fecal samples and used for
175	in vitro culture in intestinal porcine epithelial cells (IPEC-J2, ACC 701, Leibniz Institute DSMZ
176	GmbH, Braunschweig, Germany) as described previously (Worliczek et al., 2013) with
177	modifications as described here. For the production of glass cover slips with confluent layers of
178	host cells infected with C. suis, cover slips were placed into 6-well culture plates for suspension
179	cultures (PAA, Pasching, Austria). IPEC-J2 were suspended in DMEM/Ham's F12 medium
180	(PAA, Austria) supplemented with 5 % fetal calf serum, 2 mM L-glutamine, 100 U/ml penicillin
181	and 0.1 mg/ml streptomycin, and seeded in a density of $4x10^5$ cells/well. After overnight
182	incubation at 37°C, 5 % CO ₂ cover slips with attached IPEC-J2 were transferred to 6-well
183	surface-treated culture plates for adherent cells (PAA, Austria). Sporozoites of C. suis were
184	excysted as described previously (Worliczek et al., 2013), suspended in culture medium and
185	applied to the host cells in a density of sporozoites:host cells of 1:30. The infected cell cultures
186	were incubated at 37°C, 5% $\rm CO_2$ and culture medium was exchanged at 1, 4, and 7 days post
187	infection (dpi).

188

189 2.3 S. neurona culture and immunofluorescence assays

S. neurona strain SN3 was cultured, transfected and processed for immunofluorescence as
described before (Dubey et al., 2017; Howe et al., 2018). In short, parasites were cultured and
maintained in bovine turbinate (BT) cell monolayers. For IFA, extracellular merozoites were
used to infect BT cells in 24-well plates containing coverslips. Typically, on day 3 post infection,
infected BT cell monolayers were methanol-fixed for downstream immunofluorescence
experiments. Freshly isolated extracellular *S. neurona* merozoites were used for transient
expression of YFP-tagged TgIMC15 (Anderson-White et al., 2011).

197

198 2.4 *B. bigemina* immunofluorescence assays

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199 B. bigemina indirect immunofluorescence assays were performed on >10% parasitemia bovine 200 red blood cells by air drying drops in 10-well slides (Electron Microscopy Sciences), followed 201 by 100% methanol fixation for 5 min at -20°C, 3 rinses in PBS and blocking in 3% BSA in PBS 202 for 1 hr at RT. Primary antibodies diluted in blocking solution (rabbit TgCentrin1 1:4000 (Fung 203 et al., 2012), mouse MAb 12G10 α-tubulin 1:100 (University of Iowa Hybridoma Bank) (Jerka-204 Dziadosz et al., 1995); mouse ascites MAb B-5-1-2 α-tubulin 1:200 (Sigma-Aldrich #T5168); rat 205 TgEB1 1:100 (Chen and Gubbels, 2019); guinea pig BbIMC1a 1:500) were incubated for 1 hr at 206 RT, followed by 3 washes for 5 min in PBS, Alexa488 or 594 conjugated secondary antisera 207 (ThermoFisher) diluted 1:400 in 0.1% BSA in PBS for 45 min at RT followed by 1 wash for 5 208 min in PBS containing 1.5 µg/ml 4',6-diamidino-2-phenylindole (DAPI) to stain DNA, and two 209 additional 5 min washes with PBS before mounting with Fluoro-Gel (Electron Microscopy 210 Sciences). A Zeiss Axiovert 200 M wide-field fluorescence microscope was used to collect 211 images, which were deconvolved and adjusted for phase contrast using Volocity software 212 (Quorum Technologies). SR-SIM was performed on a Zeiss ELYRA S.1 system in the Boston 213 College Imaging Core in consultation with Bret Judson. All images were acquired, analyzed and 214 adjusted using ZEN software and standard settings.

- 215
- 216

2.5 *C. suis* indirect immunofluorescence assays

217 C. suis indirect immunofluorescence assays were performed on infected IPEC-J2 cells grown on 218 glass cover slips as described above at dpi 7 and 8. Infected cells on cover slips were washed in 219 PBS, fixed with ice cold methanol for 10 min and subsequently washed twice with PBS. 220 Blocking was performed for 20 min with Superblock® T20 (PBS) blocking buffer (Thermo 221 Scientific) with 1% (v/v) normal goat serum at RT. Primary antibodies (guinea pig TgNuf2 222 (Farrell and Gubbels, 2014); mouse mAb 6-11B-1 acetylated α-tubulin, Invitrogen) were diluted 223 to 1:1000 (v/v) in blocking solution, and incubated for 90 min at 37° C, followed by 2x10 min 224 washes with PBS-Tween 20 (0.01%, v/v), incubation with secondary antibodies diluted 1:600 in 225 blocking solution for 40 min at 37°C, 2x10 min washes with PBS-Tween 20, followed by 1 wash 226 for 5 min in PBS containing 1 µM DAPI, 4 washes of 5 min at RT with PBS and quick 227 immersion in ddH₂O, before mounting with Aqua Polymount (Fisher Scientific). Mounted 228 samples were hardened for 2-5 days before imaging. A Zeiss LSM 510 Meta with a 63x plan 229 apochromat oil immersion objective was used to collect Z-stacks using ZEN 2009 Light Edition

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230 (Carl Zeiss Microimaging GmbH, Jena, Germany). Z-stacks were subsequently deconvoluted

using Huygens Essential 4.3 software (Scientific Volume Imaging Inc., The Netherlands) with

232 measured point spread functions (PSF) for the respective channels. Z-projections (maximum

233 intensity projections of split channels) were computed with ImageJ 1.48e (National Institutes of

Health, Bethesda, MD, USA).

235

236 **2.6** Transmission electron microscopy

For thin-section transmission, pellets of >10% parasitemia *B. bigemina* infected bovine red blood

cells were fixed with 2.5% glutaraldehyde in 0.1 M sodium cacodylate (Sigma) buffer (pH 7.4)

for 1 hr at RT and processed as described (Nishikawa et al., 2005) before examination with a

240 Philips CM120 Electron Microscope (Eindhoven, The Netherlands) under 80 kV.

241 Pellets of >10% parasitemia B. divergens infected human red blood cells were fixed with 2.5% 242 glutaraldehyde 1.25% paraformaldehyde and 0.03% picric acid in 0.1 M sodium cacodylate buffer (pH 7.4) overnight at 4°C. Cells were washed in 0.1 M cacodylate buffer and post fixed with 1% 243 244 OsO₄/1.5% KFeCN₆ for 1 hr, washed 2x in water, 1x maleate buffer (MB) 1x and incubated in 1% 245 uranyl acetate in MB for 1 hr followed by 2 washes in water and subsequent dehydration in grades 246 of alcohol (10 min each; 50%, 70%, 90%, 2x10 min 100%). The samples were then put in 247 propyleneoxide for 1 hr and infiltrated overnight in a 1:1 mixture of propyleneoxide and TAAB 248 (TAAB Laboratories Equipment Ltd). The following day the samples were embedded in TAAB 249 Epon and polymerized at 60°C for 48 hrs. Ultrathin ~60 nm sections) were cut on a Reichert 250 Ultracut-S microtome, picked up on to copper grids stained with lead citrate and examined in a 251 JEOL 1200EX Transmission electron microscope and images were recorded with an AMT 2k 252 CCD camera.

253

254 **2.7 Phylogenetic analysis.**

255 The following 18S rRNA gene sequences were used: Cryptosporidium parvum (GenBank

accession no.: L25642.1), Cryptosporidium muris (AB089284.1), Babesia bovis (L19077.1),

257 Babesia bigemina (JQ437261.1), Babesia divergens (AJ439713.1), Babesia duncani

258 (HQ285838.1), Babesia microti (AB219802.1), Theileria annulata (EU083801.1), Theileria

259 parva (L02366.1), Theileria orientalis (HM538200.1), Cytauxzoon felis (L19080.1), Theileria

260 equi (KM046918.1), Hepatozoon canis (KX712124.1), Plasmodium falciparum

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- 261 (XR 002966654.1), Plasmodium berghei (XR 002688202.1), Plasmodium vivax
- 262 (XR 003001225.1), Toxoplasma gondii (L24381.1), Hammondia hammondi (KT184369.1),
- 263 Neospora caninum (U03069.1), Besnoitia besnoiti (KJ746531.1), Cystoisospora suis
- 264 (KF854251.1), Sarcocystis neurona (KT184371.1), Hyaloklossia lieberkuehni (AF298623.1),
- 265 Goussia janae (GU479644.1), Goussia neglecta (FJ009242.1), Cyclospora cayetanensis
- 266 (XR_003297358.1), Eimeria callospermophili (JQ993648.1), Eimeria tenella (KT184354.1),
- 267 Eimeria necatrix (KT184349.1), Eimeria maxima (KT184346.1), Eimeria brunetti
- 268 (KT184337.1), and *Eimeria acervulina* (KT184333.1). Sequences were aligned using Clustal
- 269 Omega (Sievers et al., 2011) and a consensus phylogenetic tree was generated using Geneious
- 270 Prime V2019.1.3 (Invitrogen) using the Jukes-Cantor genetic distance model, neighbor-joining
- tree build method, bootstrapped 100x using *C. parvum* as outgroup.
- 272

273 2.8 Generation of *B. bigemima* IMC1a antiserum

- 274 We identified five *B. bigemina* IMC proteins by reciprocal BLASTP searches on EuPathDB
- 275 (Warrenfeltz et al., 2018) with the 14 T. gondii alveolin-domain containing IMC proteins
- 276 (Anderson-White et al., 2011): PiroplasmaDB.org accession number BBBOND_0204530,
- 277 BBBOND_0401990, BBBOND_0201220, BBBOND_0208040, BBBOND_0300730). B.
- 278 *bigemina* BBBOND_0204530, which we named BbIMC1a, was most conserved and the whole
- 279 ORF of 600 bp was synthesized by TwistBioscience (San Francisco, CA), amplified with
- 280 primers #4831 pAVA-Gibson-Twist-F gaagctcagacccaggg and #4832 pAVA-Gibson-Twist-R
- 281 tgcagaacttgttcgtgctg to remove the linkers and cloned by Gibson assembly into PmeI/NruI
- digested plasmid pAVA0421 (Alexandrov et al., 2004), expressed as a 6-His tag fusion in
- 283 Escherichia coli BL21-RIPL, purified by Ni-NTA chromatography under denaturing conditions
- 284 (Invitrogen), refolded, and used to immunize a guinea pig (Covance, Inc). Serum was affinity
- purified as described previously (Gubbels et al., 2006) against recombinant His6-BbIMC1a.
- 286

287 3 Results

288 **3.1 External budding**

Schizogony is defined as: schizo = split (or cleft); gony = birth (genesis of a class of thing). It is
generally understood as meaning "multi-fission" and is applied to division modes producing
more than two daughter cells per division by peripheral (or cortical) budding from the plasma

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membrane of a polyploid, multi-nucleated mother cell (Fig 2). However, confusion enters with
the term "schizont", which is more widely defined and used to describe any polyploid
intermediate cell state regardless of division mode (Fig 4-6): schizo = split (or cleft); -ont = a
being (from Greek einai to be) i.e., "to be split". We forewarn that the term 'schizont' is and will
be used across division modes for polyploid cells, and thus will not always refer to schizogony
per se.

298

299 3.1.1 Schizogony with karyokinesis

300 A wide array of Apicomplexa impacting humans or livestock divide by schizogony, such as 301 nearly all Eimeriidae family members of the Coccidian parasites (Dubremetz, 1973; Joyner and 302 Long, 1974; Dubremetz and Elsner, 1979; Ferguson et al., 2008), the genus of *Plasmodium* in the 303 order of Haemosporida (Arnot et al., 2011; Stanway et al., 2011) as well most (but not all!) 304 piroplasms compromising *Theileria* and *Babesia* spp. (Mehlhorn and Shein, 1984) (Fig 1). The 305 piroplasms are tick-transmitted parasites that multiply in red blood cells without forming 306 pigments (order Achromatorida). Both the piroplasms and the *Plasmodium* spp. (order 307 Chromatorida: form pigments in red blood cells) reside in the clade Haemosporidia for their 308 shared residence within red blood cells. Our discussion here is based mostly on Plasmodium 309 schizogony in the red blood cells as it is best studied system, however, insights likely apply 310 widely. A defining principle of schizogony is the disassembly of the mother's cytoskeleton 311 shortly following completion of host cell invasion, resulting in an amoeboid or pleomorphic cell 312 (Hepler et al., 1966; Gruring et al., 2011) (Fig 2A2) which then undergoes several cycles of 313 DNA replication and nuclear division (Fig 2A3,4). These rounds of DNA replication without 314 budding are controlled by cdc2-related kinase CRK4 which resides in the nucleoplasm and is 315 associated with phosphorylation of the DNA replication machinery (Ganter et al., 2017). 316 Interestingly, the mitotic cycles of the nuclei sharing the same cytoplasm are not synchronous. 317 Mechanistically, this observation is anchored in a differential maturation model of mother and 318 daughter centrosomes following their duplication. Although this model has been firmly 319 demonstrated in higher eukaryotes and is defined by proteins in the centrosome's distal and sub-320 distal appendages (Bornens and Gonczy, 2014), it has not been formally confirmed in the 321 Apicomplexa. The last cycle of S/M-phase, in contrast, is synchronous and coupled to the 322 synchronous budding of the daughters at the end of schizogony (Ferguson et al., 2008; Kono et

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323 al., 2016; Rudlaff et al., 2019) (Fig 2A4,5). Budding follows the activation of the centrosomes' 324 outer-cores, which is always tied to simultaneous centrosome inner-core activation. The exact 325 timing of the nuclear cycle before the onset of budding has not been clearly resolved. Notably, 326 the level of PfCRK4 drops when the budding cycle is about to start, but it is not clear whether 327 this is the defining signal synchronizing the nuclear cycles and/or activating the budding cycle. 328 Potentially, a diffusible signaling protein could respond to the depletion of nutrients, the 329 accumulation of waste, limited space, or act as a quorum sensor, and subsequently synchronize 330 or halt the nuclear cycle before activating the synchronous budding phase. Since the number of 331 offspring varies across *Plasmodium* species, even between different *P. falciparum* strains, across 332 different development stages (e.g. the liver cell expands and produces up to 90,000 merozoites 333 from infection by a single sporozoite (Vaughan and Kappe, 2017)) there is also a strong, 334 programmed genetic component determining the number of offspring (Reilly et al., 2007). Either 335 way, when the 'commitment to budding' checkpoint is cleared, the centrosomes that anchor the 336 spindle pole and chromosomes, reorient to associate with the plasma membrane, a step which is 337 mediated by Cyc1 (Robbins et al., 2017). Note that the centrosomes in *Plasmodium* spp. 338 technically are centriolar plaques as these parasites lack canonical centriolar structures. 339 Following plasma membrane docking of the centrosomes, the daughter cytoskeletons start to 340 assemble in connection with the plasma membrane and the buds move outward (Fig 2A4,5). In 341 P. falciparum the maturation and release of daughters is mediated by CINCH, a contractile 342 protein in the basal complex (Rudlaff et al., 2019).

343

344 3.1.2 Schizogony with limited karyokinesis

345 Although not separately recognized in the naming conventions, a variation on the classical 346 schizogony as described in the preceding section occurs during sporozoite formation in the 347 invertebrate vector wherein sexual development takes place (Fig 1). This happens in the 348 mosquito midgut for the *Plasmodium* spp (Simonetti, 1996) and in the tick salivary gland for 349 most piroplasms, including *Theileria* spp. and many *Babesia* spp. (Mehlhorn and Shein, 1984; 350 Jalovecka et al., 2018). P. berghei and P. falciparum sporogenesis have been most extensively 351 studied, which occurs in an extracellular oocyst residing under the basal lamina of the midgut 352 wall (Simonetti, 1996). Like in classical schizogony, the mother's cytoskeleton is disassembled 353 resulting in a pleomorphic cell. Here, DNA replication and mitosis are not always followed by

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354 karyokinesis leading to a patchwork of nuclei, with varying levels of ploidity (Fig 2B). Again, 355 the individual nuclei are at different levels of S-phase and mitosis, however, within one 356 polyploid nucleus this cycle is largely synchronous (Howells and Davies, 1971b; a; Schrevel et 357 al., 1977; Sinden and Strong, 1978). Typically, a round of sporogenesis produces thousands of 358 individual sporozoites from a single mother cell. Similar to classical schizogony, the budding of 359 daughters from the cortex occurs simultaneously and is coupled to a synchronized round of S-360 phase and mitosis (Howells and Davies, 1971a; Schrevel et al., 1977; Araki et al., 2019; Pandey 361 et al., 2019). This process is conserved in *Theileria* and *Babesia* spp. as well, where multiple and 362 multi-lobed nuclei are especially prominent (Moltmann et al., 1983). However, some Babesia 363 spp., produce sporozoites through binary fission (Mehlhorn and Shein, 1984). 364 In summary, the intriguing phenomenon is that karyokinesis seems to be optional in these

polyploid schizonts. The synchronized cycles of mitosis within a single nucleus suggest that mitotic cycles are organized on the nucleoplasm level just as in classical schizogony, whereas commitment to budding is a factor shared across the whole cytoplasm. How these events are controlled at the molecular level has not been determined.

369

370 3.1.3 Binary and multiple fission

371 Binary fission is defined as "the formation of two daughter cells per division round". Among the 372 Approximation Approximation and the second s 373 of many Babesia and Theileria parasites. Most Babesia spp. form two daughters per division 374 round, i.e. classical binary fission. However, in the red blood cell cycle several clades of 375 Babesia, including Babesia duncani (Conrad et al., 2006) and B. microti (Rudzinska, 1981), 376 form four merozoites per division round (Maltese cross), as do all Theileria spp. (Conrad et al., 377 1985; Conrad et al., 1986; Fawcett et al., 1987; Uilenberg, 2006). This process is known as both 378 schizogony and 'multiple fission', however it is generally considered distinct from schizogony, 379 which as mentioned above, is a term historically used when many more daughters are produced. 380 The term "binary fission" does not imply that budding is involved and does not intuitively 381 connect with "multiple fission", which produces only four daughters per division round, as 382 defined in schizogony. Due to the lack of clarity on how these processes are related and defined 383 cell biologically, the exact nature of Babesia cell division in the red blood cell remains to be 384 elucidated. The ambiguity of the used terms (i.e. binary fission, budding, schizogony) to describe

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385 this process further exacerbates this confusion. In a 1978 publication study using transmission 386 electron microscopy (TEM) the interpretation was as follows (Potgieter and Els, 1977): "The 387 trophozoites were surrounded by a single membrane, were pleomorphic in shape and contained 388 large inclusions of host cell cytoplasm, but no cytostomes or food vacuoles could be identified. 389 Reproduction took place through a process resembling schizogony resulting in the production of 390 two merozoites, the cytoplasmic constituents of the original trophozoite (mother cell) being 391 virtually entirely incorporated into the daughter cells in the process". In essence, the 392 pleomorphic/ameboid trophozoite in combination with the schizogony reference suggest that the 393 process is conceptually very similar to schizogony, except that term does not capture it since 394 only two daughters are formed. Here we combine existing data with new insights from *in vitro* 395 cultivated Babesia bigemina to clarify the mechanistic murkiness surrounding binary fission. 396 *B. bigemina* is a cattle-infecting, tick-transmitted apicomplexan causing vast economic losses 397 (Bock et al., 2004; Suarez et al., 2019). From an experimental point of view, *B. bigemina* cell 398 division is uncomplicated in that it only produces two daughters per division round, and it is a 399 relatively large *Babesia* spp., making it an ideal candidate for microscopy. We focused on the 400 key organizers of cell division: the centrosome (or centriolar plaques) and cortical cytoskeleton. 401 T. gondii Centrin1 antiserum on B. bigemina showed a specific, albeit relatively weak, signal, 402 but only during daughter budding (Fig 3A). It therefore appears that centrosome composition is 403 dynamic, which was recently also reported for P. berghei Centrin4 as this signal was only on the 404 centrosome during mitosis and budding, but not in mature or recently invaded parasites (Roques 405 et al., 2019). In contrast, *Plasmodium* Centrin3 is a popular marker of centrosomes and is always 406 associated with the centrosome, indicating that the dynamics of different Centrins can vary 407 (Mahajan et al., 2008; Kono et al., 2012). We detected microtubules with α -tubulin MAb 12G10 408 generated against ciliate Tetrahymena thermophila (Jerka-Dziadosz and Frankel, 1995), which 409 has shown broad reactivity in the Apicomplexa. A spot focused toward the apical end was 410 observed (Fig 3A), but we never observed tubulin in the nucleus, suggesting this antibody does 411 not detect the mitotic spindle. In *Plasmodium* the spindle was successfully visualized with 412 monoclonal antibody B-5-2-1 generated against sea urchin α-tubulin (Gerald et al., 2011), but in 413 B. bigemina we observed the same patterns as for MAb 12G10 (not shown). In T. gondii, 414 detecting the spindle microtubules has also been challenging, but a polyclonal serum against 415 TgEB1, a microtubule (+)-end binding protein, was very specific for the spindle (Chen et al.,

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416 2015b). Reactivity of α -TgEB1 in *B. bigemina* across all parasite stages highlighted a spot basal 417 of the apical tubulin staining (Fig 3B). We interpret that EB1 does not bind to the spindle 418 microtubules in *B. bigemina*, but instead associates with the (+)-end of the subpellicular 419 microtubules emanating from the MTOC at the very apical end. This also indicates these 420 microtubules are relatively short and only cover the very apical cap of the parasite. To better 421 visualize the cortical cytoskeletal scaffolds that drive budding we tested polyclonal antisera 422 generated against T. gondii IMC proteins (Gubbels et al., 2004; Anderson-White et al., 2011), 423 but unfortunately none showed a signal. Therefore, we generated a specific B. bigemina IMC 424 antiserum against BBBOND 0204530, the closest relative to the *T. gondii* alveolin domain 425 containing IMC proteins (Anderson-White et al., 2011), which we named BbIMC1a (Fig S1). 426 IFAs with α -PbIMC1a nicely highlighted the cortical cytoskeleton during budding, extending 427 basally beyond the microtubule signal. In mature parasites we observed signal along the entire 428 length of the merozoite (Fig 3C). In trophozoites we also observed parasites identified by their 429 DAPI signal with weak or variable intensities of either the IMC1a or tubulin signals that 430 appeared disorganized, or at least not to outline a merozoite (Fig 3C top panel and middle panel 431 marked by an asterisk). We interpreted these amorphic signals to represent trophozoites that are 432 at the stage between invasion and the start of cell division. However, across all parasites in 433 division we observed parasites forming next to each other, with apical ends emerging first with 434 V-shape symmetry.

435 To obtain a higher resolution of the mechanistic steps in the *B. bigemina* division process we 436 performed TEM. We focused on stages after the parasites escaped from their vacuole and entered 437 G1 growth phase (trophozoites). We observed a high frequency of zoites wherein the IMC 438 breaks into 5-6 pieces, still in curved structures on the edge consistent with an ameboid or 439 pleomorphic status for the zoites (Fig 3D1). This consistent observation suggests that 440 disassembly of the mother cytoskeleton is spatially organized resulting in a symmetrical 441 appearance, which reflect the trophozoites identified by IFA (Fig 3C). Similar observation on 442 disassembling mother IMC have been reported for *Plasmodium* ookinetes in the mosquito 443 midgut (Carter et al., 2007), and for sporozoites in liver cells, where regularly spaced breaks 444 occur in the IMC, although not quite as symmetrically organized as seen here in *B. bigemina* 445 (Jayabalasingham et al., 2010). In a budding zoite two daughter buds are emerging on one side of 446 the cell (like rabbit ears), and not on polar opposites (Fig 3D2). The shape of the nucleus is also

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447 consistent with the anchoring of the nucleus at the apical end of the daughters, which, by using 448 other division modes as a guide, we assume is through the spindle poles (Potgieter and Els, 449 1977). For the growing stages, we captured many transverse cross sections of two side-by-side 450 budding daughters with the nucleus at various stages of division (Fig 3D 3-5). Finally, 451 completely formed daughters appear side by side under a similar angle as the early daughter buds 452 (compare the orange arrows in Fig 3D2 with 3D6). Additional TEM images of *Babesia* buds in a 453 V orientation consistent with our data have been reported elsewhere (Friedhoff and Scholtyseck, 454 1977; Potgieter and Els, 1977; Scholtyseck, 1979). A section through the basal ends of still 455 connected *Babesia divergens* parasites (predominantly a cattle parasite but opportunistic in 456 humans) displays an electron dense structure on the basal end of the IMC consistent with a basal 457 complex (Fig 3E). Although we did not capture a section through this structure in *B. bigemina*, a 458 similar appearance of the basal complex in *B. bigemina* has been reported previously (Potgieter 459 and Els, 1977).

460 Overall, the budding of daughters in *B. bigemina* shares many features seen across asexual 461 apicomplexan development (Fig 3F). First, the angle between the daughter buds fits with the 462 pleuromitosis (closed mitosis with the spindle poles in close proximity at one side of the nucleus) 463 model dictated by closely apposed centrosomes at the apically defining side of the nucleus. This 464 is observed across the Apicomplexa that have been studied (Gerald et al., 2011; Francia and 465 Striepen, 2014). As a result, daughter buds are formed next to each other rather than in opposing 466 orientations (middle panel Fig 3F3-5). The *B. bigemina* trophozoite has an ameboid appearance 467 after invasion and cytoskeleton is disassembled (Fig 3E1, 3F2; and (Potgieter and Els, 1977)), 468 which likewise is observed during the corresponding stage of *P. falciparum* (Gruring et al., 469 2011). Taken together, these data firmly illustrate that the processes known as binary fission and 470 schizogony, and by extension multiple fission as seen in *Theileria* spp., are mechanistically the 471 same and only differ in the number of nuclear replication cycles before the onset of cytokinesis. 472 Collectively, we now group these division forms together under the umbrella "external (aka 473 cortical) budding"

474

475 **3.1.3** Other variations on cortical budding

Schizogony by cortical budding as described in the preceding sections are found in variousasexual stages. Other forms on cortical budding can be found in sexual development stages,

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478 which we will briefly highlight here to illustrate the spectrum of division modes. *Plasmodium* 479 male gametocytogenesis produces flagellated microgametes in a process known as 480 "exflagellation". This process occurs in the red blood cell and unfolds by several fast rounds of 481 S/M-phase resulting in a single polyploid nucleus (Sinden et al., 1978; Sinden, 1983). The 482 cortical 'budding' of microgametes, which at the cytoskeletal level basically are a single 483 flagellum and only contain microtubules, progresses while the nuclear material partitions. 484 Interestingly, a recent study using Ndc80 as a marker for the kinetochores demonstrated that the 485 genome size universally increases to 8N, but that chromosome replication is asynchronous 486 (Pandey et al., 2019). This is in contrast to what is observed during asexual schizogony, where 487 the rule is that the mitotic cycles are synchronous within a shared nucleoplasm. 488 In addition, microgametocyte formation of T. gondii also progresses through a cortical budding process (Ferguson et al., 1974). As discussed in detail below, it is salient to note that the 489 490 asexual division of T. gondii is by internal budding, not through cortical budding. T. gondii 491 microgametocytes formation plays out by the association of 1N nuclei in a multi-nucleated 492 microgamont with plasma membrane of the mother cell. Interestingly, the mother cell 493 (macrogamont) maintains an IMC where in holes are present to facilitate association of the 494 nuclei with the plasma membrane. From here, bi-flagellated microgametes bud outward in a 495 cortical budding process (Ferguson et al., 2008). Interestingly, this process is conserved across 496 the Coccidia, as it is also reported for *Eimeria* spp. which, in contrast to *T. gondii*, replicate 497 asexually by schizogony (Ferguson et al., 1980) (Fig 1). Taken together, the notable features here 498 are maintenance of the cortical cytoskeleton (Dubey et al., 2017) in combination with cortical 499 budding while S-phase, mitosis and karyokinesis have been completed before the onset 500 'budding'.

501

502 **3.2** Internal budding

503 3.2.1 Endodyogeny

Endodyogeny is defined as: endo = inner/internal; dyo = two; geny = genesis, meaning birth;
production/generation/origin. This term is applied to a type of reproduction in which two
daughters are formed within a parent cell (Goldman et al., 1958; Sheffield and Melton, 1968).
Endodyogeny has been described in detail for *T. gondii* tachyzoites (Nishi et al., 2008;

508 Anderson-White et al., 2012) and is employed as well by its closest evolutionary neighbors

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509 comprising several genera of the Sarcocystidae (Fig 1, 4). These include Hammondia, Neospora 510 and *Besnotia* spp. as well as the tissue cyst-forming stages of *Sarcocystis neurona*. During 511 endodyogeny, the mother's cytoskeleton is not disassembled following completion of invasion, 512 and two daughter buds assemble on centrosomes residing within the cytoplasm. Only at the very 513 last stage of daughter budding is the mother's cytoskeleton disassembled and is the plasma 514 membrane deposited on the new daughters, which is mediated by recruitment of the "gliding 515 motor complex" to the IMC. This complex contains a multi-acylated protein glideosome 516 associated protein, GAP45, that is anchored in both the plasma membrane and the IMC outer 517 membrane and 'zippers' these structures together in an apical to basal direction (Gaskins et al., 518 2004; Frenal et al., 2014). Many of the cell cycle checkpoints throughout endodyogeny have 519 been resolved. Specifically, two checkpoints have been described upon commitment to mitosis 520 and budding, one likely dedicated to mitosis and the other to budding. This is thought to facilitate 521 the uncoupling of S/M cycles from budding in the multi-daughter division modes and 522 differentially activate the centrosome inner- and outer-cores (Suvorova et al., 2015; Naumov et 523 al., 2017; White and Suvorova, 2018). Like in schizogony, daughter budding occurs in sync with 524 S/M-phase and karyokinesis. Upon emergence of daughters, a narrow cytoplasmic bridge at the 525 basal end remains connected with a residual body containing remnants of the mother cell, which 526 will be largely resorbed into the daughters (Frenal et al., 2017b; Periz et al., 2019). As a result, 527 the daughter parasites remain in contact with each other and could therefore have the false 528 appearance of endopolygeny. To differentiate these two processes, this division process has been 529 referred to as 'repeated endodyogeny'.

530

531 3.2.3 Endopolygeny

532 Endopolygeny is defined as: endo = inner/internal; poly = multiple; geny= genesis/birth. This 533 term is used to describe the reproduction types wherein more than two individuals are formed 534 simultaneously within the cytoplasm (i.e. not from the cortical periphery) of a polyploid parent 535 cell. One of the key features of internal budding is that the mother's cytoskeleton is maintained 536 throughout cell division and only dissembles just before the emergence of almost completely 537 assembled daughters. Although not differentiated in the naming conventions, two sub-forms can 538 be distinguished: either the polyploid mother cell can be multinucleate, or it can contain one 539 large polyploid nucleus depending on whether karyokinesis follows each round of S-phase. Both

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540 these forms of budding are found within the tissue cyst-forming Coccidia, more specifically the 541 Sarcocystidae (Fig 1, 5, 6). These division forms are related to endodyogeny, since the mother's 542 cytoskeleton is maintained throughout the division cycle until the maturation of the daughter 543 cells. We define these three forms here collectively as "internal budding". Internal budding by 544 the Sarcocystidae is unique among the Coccidia as other families, notably the closely related 545 Eimeriidae, replicate by schizogony (Fig 1). Two emerging systems exist to study the variations 546 in progression of endopolygeny: Sarcocystis neurona is a model for studies representing the 547 form without completing karyokinesis (Fig 5) (Vaishnava et al., 2005; Dubey et al., 2017), 548 whereas merogony of *Cystoisospora suis* provides an accessible *in vitro* model for the form 549 including karyokinesis after each S/M-round (Fig 6) (Worliczek et al., 2013).

550

551 **3.2.3.1 Endopolygeny with karyokinesis**

552 C. suis is the causative agent of suckling piglet coccidiosis (Shrestha et al., 2015). Although

553 phylogenetically *C. suis* falls within the cyst-forming Coccidia (Fig 1), tissue cysts have never

been observed and the known coccidian development cycle is monoxenic in pigs: the absence of

tissue cyst forming capacity is likely a secondary loss in this species (Stuart et al., 1982; Shrestha

et al., 2015). *C. suis* asexually replicates in epithelial cells of the porcine small intestine by

557 endodyogeny and endopolygeny, depending on the generation of asexual stages (alternatively

called types: meronts/merozoites type I, II and subtype II (Matuschka and Heydorn, 1980). Early

asexual division after infection of the gut epithelium is restricted to endodyogeny (often in

560 consecutive cycles within one parasitophorous vacuole, i.e. 'repeated endodyogeny'), whereas

561 endopolygeny is described from day 3-4 post infection onwards *in vivo* (Lindsay et al., 1980;

562 Matuschka and Heydorn, 1980) and from day 7 onward *in vitro* (Worliczek et al., 2013),

563 concurrent with cells replicating by endodyogeny.

As for *B. bigemina*, we tracked *in vitro* progression of *C. suis* development using the toolbox of reagents we established for *T. gondii*. In *C. suis* zoites undergoing endopolygeny, the mother's subpellicular microtubule cytoskeleton is clearly visible in large polyploid cells as an apically concentrated microtubular accumulation joined in the mother's conoid (Fig 5A-D). This confirms the mother's cytoskeleton is maintained throughout endopolygeny. To track the progression of S/M-phase across the nuclei we used the kinetochore component Nuf2 (Farrell and Gubbels, 2014) in combination with acetylated α -tubulin, which marks the spindle poles

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during mitosis (note that spindle microtubules disassemble during interphase (Farrell and 571 572 Gubbels, 2014; Chen et al., 2015b)). The first observation is that the status of mitosis (spindle 573 visible by tubulin stain) and the status of kinetochore separation varies among nuclei within the 574 same cell (Fig 5A, B). Furthermore, it is evident that parasites expand non-geometrically, as 575 counted by spindle poles per nucleus varying from 3, 4, 5, to 6 are seen in Fig 5A and B, whereas 576 12 budding daughters, i.e. non-geometrically expanded, can be discerned in Fig 5C, D. These 577 two observations indicate that the nuclear division cycles are not synchronous, which is 578 consistent with expansion numbers described in the literature (e.g. (Lindsay et al., 1980; 579 Matuschka and Heydorn, 1980)). Here we show for the first time that this is associated with 580 asynchronous nuclear replication cycles. Finally, we show that the final round of mitosis is 581 synchronous for all nuclei and is coupled with daughter budding, as seen in schizogony. Our 582 insights are summarized in the schematic of Fig 5E. Based on the details revealed here, we 583 conclude that like in schizogony, the S/M-phase progression are controlled at the nuclear level, 584 most likely differing maturity of the mother and daughter centrosomes, but that the commitment 585 to budding is synchronized across the cytoplasm. 586 Reports on T. gondii endopolygeny suggest geometric expansion of nuclei: 8–16 progeny per

endopolygeny replication cycle in the cat gut have been observed (Ferguson et al., 1974).
Historically, the sexual cycle of *T. gondii* has been poorly experimentally accessible, but *in vitro*completion of the sexual cycle in cat intestinal organoids was recently reported, which is
expected to provide experimental accessibility (Martorelli Di Genova et al., 2019). To date, *T. gondii* endopolygeny studies have not been so comprehensive to support a strong conclusion in
this matter.

593 An additional notable observation in Fig 5A is the presence of multiple parasites within a 594 single vacuole undergoing endopolygeny. It has been described before that several multinucleate 595 meronts can be present in a single vacuole, which indicates that more than one asexual division 596 occurs in the same cell (Lindsay et al., 1980). As such, this reveals that multiple rounds of 597 budding within the same vacuole are not exclusively found during endodyogeny as described 598 above for *T. gondii*. This phenomenon therefore highlights that the timing of budding is 599 determined by a genetic program, and not controlled by environmental factors, which we also 600 concluded for schizogony.

601

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602 **3.2.3.1 Endopolygeny without karyokinesis**

603 S. neurona has a sylvatic cycle in the Americas with small mammals as intermediate hosts and 604 the opossum as the definitive host. However, accidental infection of horses can cause equine 605 protozoal myeloencephalitis (EPM) (Reed et al., 2016). Using the same strategy as mentioned 606 above, we observe an apical concentration of microtubules across cells with progressively larger 607 single nuclei indicating the mother's cytoskeleton is maintained throughout endopolygeny 608 (arrowheads in Fig 6A). Furthermore, the S. neurona IMC can be visualized through 609 overexpression of TgIMC15-YFP, which highlights the sutures in the IMC of the mother 610 throughout development (Fig 6B) (Dubey et al., 2016). When the cell prepares for budding 611 IMC15 also appears on the centrosomes at which point the mother's IMC is still prominently 612 present (Fig 6B2). Only at the conclusion of the budding process is the mother's cytoskeleton 613 disassembled, which mimics the dynamics of the IMC during T. gondii endodyogeny (Anderson-614 White et al., 2011; Dubey et al., 2016). 615 In contrast, to C. suis, the S/M cycles of S. neurona are completely synchronized: in Fig 6A2 616 we observe a cell in interphase characterized by absence of spindle microtubules, whereas the 617 right cell displays spindle microtubules associated with each centrosome. This confirms previous 618 observations (Vaishnava et al., 2005), and indicates that when the nucleoplasm is shared, the 619 individual status of centrosome maturation is overruled, likely by a factor diffusing in the 620 nucleoplasm. Late in development the large polyploid 32N nucleus shifts into a multi-lobed. 621 'serpentine' morphology (Fig 6A3), which sets the stage for a final round of S/M-phase now 622 coupled to karyokinesis and internal budding resulting in 64 daughters (Fig 6A4). Thus, when 623 the S/M-phase are not followed by karyokinesis the nuclear cycles remain synchronized and 624 progeny number are the result of geometric expansion.

625

626 **3.2.4** Budding without multiplication and multiplication without budding

The spectrum of division modes not yet covered spans two more manifestations in sexual
development stages. The first, budding without multiplication, is the formation of kinetes in the
Heamosporidia. The *Plasmodium* spp. generate ookinetes from a fertilized zygote (ploidity of 24N in a single nucleus) that cross the mosquito gut wall (reviewed in (Angrisano et al., 2012;
Bennink et al., 2016)). Although the zygote lacks a cortical cytoskeleton, the ookinete has a

632 complete cortical cytoskeleton comprising IMC and cortical microtubules. This cortical

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cytoskeleton is formed by the ookinete through from the cortex of the zygote in absence of
karyokinesis, though a budding stage dubbed 'retort' (Canning and Sinden, 1973; Carter et al.,
2007). Yet even more exotic variations occur in the piroplasms; both *Theileria* and *Babesia* spp.
produce kinetes, which do not bud from the cortex, but into an internal vacuole formed inside the
zygote (see (Mehlhorn and Shein, 1984) for a detailed review). Thus, cortical budding can be
uncoupled from the nuclear cycle in the zygote stages.

639 The second variation is multiplication without budding, of which there are three different, 640 possibly related examples. The first is found in the *Babesia* spp. kinetes, which following their 641 formation in the tick gut, cross the midgut and migrate to the tick ovary. Here, the cortical 642 cytoskeleton disassembles and the cell transforms into a pleomorphic cell (Moltmann et al., 643 1982; Mehlhorn and Shein, 1984). Subsequently, several rounds of S/M-phase without 644 karyokinesis occur to produce a multiploid, lobed nucleus. In a process not understood at neither 645 the mechanistic nor molecular level, these large cells divide into multiple cells, each with a 646 single nucleus. Surprisingly, neither the mother nor the daughter cells have a cortical 647 cytoskeleton. This indicates that at this life stage cell division is independent of any form of 648 budding, and thus indicates that in Apicomplexa the cortical cytoskeleton is not a strict 649 requirement for cell division. From each of these pleomorphic cells a new kinete then forms by 650 budding mediated by cortical cytoskeleton formation into an internal vacuole as described above.

The released kinetes then migrate to the salivary glands where they undergo sporogenesis.

652 The second example of multiplication without budding is a binary variation on the above 653 process found in some *Babesia* spp. (Mehlhorn and Shein, 1984). Kinetes that invaded the tick 654 salivary gland disassemble their cytoskeleton and undergo one round of S/M-phase, karyokinesis 655 and cell division. However, no cortical cytoskeleton is assembled during these binary division 656 rounds. The cortical cytoskeleton is reportedly assembled slowly during the last couple of 657 division rounds resulting in mature sporozoites. Thus, the salient details are that this process is 658 binary and appears to be efficient with resources as the cytoskeleton is only assembled in the last 659 round (unlike endodyogeny, where a complete parasite is assembled each multiplication round). 660 This process has been described for *B. canis* (Schein et al., 1979), a dog parasite, and *B. bovis* 661 (Potgieter and Els, 1976), a cattle parasite, but likely occurs widely across *Babesia* spp. 662 exclusively replicating by binary fission in the red blood cell (Fig 1) (Mehlhorn and Shein, 663 1984). Thus, across all life stages, these particular *Babesia* parasites seem to have lost the ability

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664 to produce more than two daughters per division round, providing a model system to unravel the 665 specifics of the genetic program and regulatory network.

666 The third example is the process of sporoblast formation in the Coccidia. This occurs within 667 the oocysts released into the environment. Depending on the species, the zygotes can divide 668 themselves into 2 to 4 sporoblasts (this number is a defining feature in diagnosis of fecal oocysts; 669 not dividing is an option as well (Gardiner et al., 1998)). Inside the sporoblasts 2-4 sporozoites 670 form through a budding mechanism coupled to DNA replication. Although studied in many 671 parasites, T. gondii produces two sporoblasts and four sporozoites per sporoblast and as such is 672 the simplest form. Consistent with EM data (Ferguson et al., 1979a), it was recently shown that 673 neither microtubules nor IMC proteins were involved in T. gondii sporoblast formation (Dubey 674 et al., 2016). Considering the process and life stage, this process therefore appears akin to kinete 675 multiplication in *Babesia* spp., as described above. This thus seems to connect these two 676 parasites across a large phylogenetic distance, which begs the question whether this is an 677 ancestral connection, or a case of convergent evolution.

678

679 **4. Discussion**

680 The picture of cell division across all life stages of the Apicomplexa is that principle differences 681 exist between the sexual and asexual cell division strategies. In the sexual stages, cell number 682 expansion is not necessarily related to budding a daughter cytoskeleton. This shows that budding 683 is in principle not required for apicomplexan cell division, but there are very few mechanistical 684 or molecular details available for these division modes. The other insight is that all invasive 685 zoites with an apical complex form by a budding strategy, which is initiated and coordinated by 686 the centrosome's outer-core and proceeds in an apical to basal assembly direction. In all the 687 asexual life cycle stages zoite budding is coupled to a complete nuclear cycle (S/M-phase plus 688 karyokinesis). Another general rule across asexual development is that budding is synchronized 689 across the whole cell. This indicates that the centrosome outer-core activation is always coupled 690 to inner-core activation in asexual stages on a shared cytoplasm-wide level (e.g. a diffusible 691 kinase or kinase substrate). The exceptions to this rule are only found in sexual stages: 692 microgametocytogenesis and (oo)kinete formation in the Haemosproridia. 693 Besides these general rules, there are several variations within asexual division modes which

694 partition into two mechanistically different strategies: internal budding and external budding (Fig

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695 1). Internal budding comprises endodyogeny and the two variations on endopolygeny, whereas 696 external budding captures the two variations on schizogony, binary fission, and multiple fission. 697 The number of offspring in each strategy can vary from two (endodyogeny and binary fission) to 698 several orders of magnitude higher (>10,000 in schizogony). At the furthest extreme are several 699 bovine-infecting Theileria spp. of which the schizonts in the white bloods that trigger 700 transformation of their lymphocyte host cells (i.e. leukemia) resulting in division and expansion 701 of the parasites schizont stage along with their host cell (Luder et al., 2009; Chakraborty et al., 702 2017). As remarked throughout, the number of daughter cells per division round in each life 703 stage is largely genetically controlled, but the details on the controls are just starting to emerge. 704 Overlapping with the genetic switch committing to budding is the bi-partite centrosome 705 cycle. The apicomplexan centrosome and mitotic cycles are controlled by Nek and Aurora 706 kinases (Reininger et al., 2011; Carvalho et al., 2013; Chen and Gubbels, 2013; Berry et al., 707 2016; Berry et al., 2018). In T. gondii it has been demonstrated that the switch from solely a 708 nuclear cycle to a combined nuclear and budding cycle is controlled by a MAP kinase-like 709 protein (Brown et al., 2014; Sugi et al., 2015; Suvorova et al., 2015). Ultimately, cell cycle 710 progression and the activation of each core is regulated by cyclin and CDK pairs adapted to each 711 apicomplexan cycle, as they likely act independently on the inner- and outer-centrosome cores 712 (Le Roch et al., 2000; Merckx et al., 2003; Alvarez and Suvorova, 2017; Ganter et al., 2017; 713 Naumov et al., 2017; Robbins et al., 2017; White and Suvorova, 2018). An open question is the 714 identity on the factor(s) controlling the pause of nuclear cycles across parasites prior to the final 715 round of coupled S/M-phase and budding in the polynucleate division modes.

716 The control of the nuclear cycle appears to depend on the ploidy of the nucleoplasm. When 717 each S/M-phase is followed by karyokinesis, then the cycles of individual nuclei are diverging. 718 Mechanistically, this observation is anchored in differential maturation of mother and daughter 719 centrosomes following their duplication; the mother centrosome is sooner primed for another 720 round of replication. This mechanism has been firmly demonstrated in higher eukaryotes and is 721 defined by proteins in the centrosome's distal and sub-distal appendages (Bornens and Gonczy, 722 2014). However, this has not been directly demonstrated yet for the apicomplexan centrosome 723 (Morlon-Guyot et al., 2017; Courjol and Gissot, 2018; Chen and Gubbels, 2019). At first sight, 724 contrasting insight comes from synchronized endodyogeny observed in T. gondii tachyzoites. 725 Upon completion of cell division, T. gondii daughters stay connected through a cytoplasmic actin

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726 bridge maintained by Myosin I (Frenal et al., 2017b; Periz et al., 2017). If this bridge is intact, all 727 conjoined tachyzoites undergo endodyogeny in synchrony resulting in geometric expansion 728 numbers. However, the cell division cycles of parasites sharing the same vacuole become 729 uncoordinated if the bridge is disrupted. Strengthening this conclusion is the observation that 4-730 5% of tachyzoites by chance form multiple (3-4) daughters per division round: the timing of 731 daughter budding is still synchronized with the other parasites in the vacuole despite the fact that 732 these tachyzoites have undergone two rounds of S/M phase (Hu et al., 2004). For signaling 733 purposes, these paradoxical "multi-daughter endodyogenic" parasites are still in the 734 synchronized cycle of "S/M coupled to budding" state consistent with endodvogeny. 735 Accumulating insights from various mutants displaying increased incidences of "multi-daughter 736 endodyogenic" parasites (Dubey et al., 2017) suggests that such parasites fail to start budding 737 because they are missing membrane building blocks to assemble the daughter IMC, and then slip 738 into the next cell cycle while remaining in the "S/M coupled to budding" state. 739 When karyokinesis does not follow each S/M phase polyploid nuclei are formed. In S. 740 *neurona* endopolygeny, karyokinesis does not occur at all till the onset of budding, but in 741 Heamosporidioan sporogenesis, karyokinesis is more optional, which leads to a mix of nuclei 742 with various levels of ploidy. However, the mitotic cycle within each nucleoplasm appears to be coordinated (Gerald et al., 2011; Roques et al., 2015), suggesting this level of control is not at the 743 744 centrosome level, but likely the chromatin or nucleoplasm level. The control and mechanism of 745 karyokinesis are not understood, which is also the case in well-studied model eukaryotes 746 dividing by closed mitosis such as *Aspergillus nidulans*, fission yeast, and baker's yeast. The 747 challenge in polyploid nuclei is to keep the multiple sets of chromosomes together so they can be

accurately partitioned into the daughters. The solution is that the chromosomes remain clustered

throughout the cell cycle by tethering the centromeres to the nuclear lamina. For instance, in *T*.

750 *gondii* the 13 centromeres and associated kinetochores remain clustered at the centrocone, a

nuclear envelope fold that houses the spindle microtubules during mitosis (Brooks et al., 2011;

Farrell and Gubbels, 2014). Similar observations have been made throughout the *Plasmodium*

⁷⁵³ life cycle using kinetochore markers (Pandey et al., 2019). Strayed chromosomes are rarely

observed but individual centromeres are seen during interphase in the *Plasmodium* schizogony in

the red blood cell. However, all centromeres cluster together again at the spindle pole before

entering mitosis (Hoeijmakers et al., 2012). Clearly, centromere clustering and sequestration at

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the nuclear envelope through the kinetochores is an effective strategy to warrant that completesets of chromosomes are maintained in polyploid nuclei.

759 None of the above addresses why there are two different forms of budding; i.e. internal vs 760 external. We will try to address this by looking at the phylogeny. The first question is whether 761 either internal budding or external budding are an innovation or ancestral process. Since not 762 enough cell division details are known for Apicomplexa outside those groups included in Fig 1, a 763 firm answer is not possible. In favor of loss of internal budding is the generally more reduced genome and streamlined biology found in the Plasmodium spp. and the piroplasms (e.g. host cell 764 765 invasion (Gubbels and Duraisingh, 2012)). Alternatively, in favor of innovation of internal 766 budding in the cyst-forming Coccidia is the putative advantage evidenced during T. gondii 767 endodyogeny. In this case, the mother parasite remains invasion- and egress-competent 768 throughout most of the replication cycle, hence increasing the resilience of the parasite in the 769 dynamic host system (i.e. immune attacks) to move between host cells (Gaji et al., 2011). In 770 contrast, during S. neurona endopolygeny the micronemes disappear halfway through the 771 division process, which depletes their invasion capacity (Vaishnava et al., 2005); hence, this 772 model does not hold true for the polyploid internal budding modes. Alternatively, it can be 773 argued that the maintenance of a cortical cytoskeleton in the mother makes the parasites more 774 resistant to mechanical stress, but it is not immediately obvious how this sets the tissue cyst-775 forming Coccidia apart from parasites dividing by external budding. As all of the Coccidia have 776 oral transmission routes, each species exhibits asexual stages in gut epithelial cells; indeed, the 777 non-cyst forming Coccidia complete their whole cycle in the intestinal epithelium. This 778 revelation suggests that the differentiation of internal from external budding might be coupled to 779 the innovation of tissue cyst formation. The requirements for cyst formation comprise dispersion 780 throughout the host beyond the gut-epithelium and assembly of the cyst in neuronal and/or 781 muscular tissue, which do not have obvious links to division modes. Both the tissue cyst 782 inhabiting bradyzoite stages of T. gondii and S. neurona (pre-bradyzoite metrocytes as well) 783 undergo endodyogeny; only the acute stage of S. neurona divides by endopolygeny (Dubey et 784 al., 2001). In addition, although the acute stage of S. neurona dissolves the vacuolar 785 compartment and replicates in the host cell cytoplasm, bradyzoite multiplication and cyst 786 formation occur within a vacuole (Jakel et al., 2001). Cyst expansion has to occur gradually so as 787 not to compromise the cyst wall, which obviously is more compatible with endodyogeny than

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endopolygeny. Notably, bradyzoite replication within *T. gondii* cysts is asynchronous and is
consistent with this model (Watts et al., 2015).

790 An outlier is *Eimeria callospermophili*, a rodent parasite that replicates by endopolygeny 791 rather than schizogony, and as far as known, does not form tissue cysts (Roberts et al., 1970; 792 Hammond, 1973). E. callospermophili undergoes several rounds of S-phase that are each 793 followed by karyokinesis to produce 4-10 nuclei per schizont while the mother's cytoskeleton is 794 maintained. In synchrony, two daughters per nucleus bud internally and the plasma membrane is 795 acquired when the daughter are about 1/3 developed, resulting in 8-20 merozoites. This suggests 796 internal budding was likely present before the advent of tissue cyst formation. However, the 797 phylogenetic position of this parasite is on the edge of the *Eimeria* spp. (Fig 1). To complicate 798 matters further, genome information has demonstrated that the *Eimeria* spp. and the *Isopora* spp. 799 branch within each other and that neither are monophyletic clades (Kvicerova and Hypsa, 2013). 800 Furthermore, the mechanism of cortical budding occurs across the Coccidia sexual cycle during 801 microgametocyte formation (Ferguson et al., 1974; Ferguson et al., 2008). Hence, parasites can 802 exhibit both cell division modes at different stages of their life-cycles – internal and external 803 budding. We also know that sporozoite formation in T. gondii occurs through an internal budding 804 mechanism in the absence of a maternal cytoskeleton (Ferguson et al., 1979b; Dubey et al., 805 2017). Interestingly, the timing of plasma membrane association with the cytoskeleton is much 806 earlier in this situation compared to the other asexual stages occurring in presence of a maternal 807 cytoskeleton, which indicates the timing of events is flexible. Taken together, there appears to be 808 a continuum between the morphogenic features of the Coccidia, suggesting that there is a 809 significant level of plasticity in biology and division modes. Further insights may result from the 810 investigation of parasite species straddling the two division modes; for example, the Goussia 811 spp. bridging the cyst-forming and non-cyst forming Coccidia (Fig 1) (Barta et al., 2001). The 812 Goussia spp. are found in fish, reptiles and amphibians, which have not been extensively studied 813 at the cell biological level (Rosenthal et al., 2016). Overall, why the two different extant forms of 814 budding exist cannot be answered with satisfaction based on the factors we considered. 815 This leaves the question of what the mechanistic differences are between internal and 816 external budding. The key question here is how parasites transitioned between the general

817 strategies of external vs internal budding. A principal difference is rooted in the localization of

the centrosome. In cortical budding the centrosomes reorient to the plasma membrane, whereas

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819 during internal budding they remain in the cytoplasm. A simple model suggests that the presence 820 of the mother's cytoskeleton physically prevents access of the centrosome to the plasma 821 membrane. Yet, it is unlikely such a simplistic model fully describes this complicated process. A 822 structure anchoring the centrosome to the cortical cytoskeleton has been described for T. gondii 823 endodyogeny; a striated fiber assemblin (SFA), extending from the centrosome to the conoid 824 (Francia et al., 2012). SFA fibers are typically found in the flagellar assembly of algae, where 825 they contribute to orientation of the basal body rootlet system relative to other subcellular 826 structures (Francia et al., 2015). During *Eimeria necatrix* schizogony, a dense structure anchors 827 the centrosome to the plasma membrane. This structure migrates basally along with the cortical 828 microtubules during the progression of daughter bud assembly (Dubremetz, 1975), and might be 829 related to the SFA fiber. Such a structure has not been described for *Plasmodium* schizogony in 830 the red blood cell, although the SFA genes are conserved in *Plasmodium* spp. (Lechtreck, 2003). 831 Thus, SFA-like structures anchoring the centrosome to the plasma membrane have been 832 observed in *Eimeria* schizogony (Dubremetz, 1971) and even in *Theileria equi* sporozoite 833 schizogony (Moltmann et al., 1983) and appear to be a differentiating factor between internal and 834 external budding. Further study of SFA genes across the Apicomplexa may reveal the nature of 835 how nuclei are anchored in the buds across the different division modes. 836 Several more principle difference between internal budding relative to cortical budding exist. 837 For example, the mother and daughter cytoskeletons must be differentially stabilized upon 838 completion of budding. What are the putative mechanisms? It has been shown that a timely 839 regulated proteolysis which removes the C-terminus of the major network component, IMC1,

840 coinciding with conversion of the network from a detergent-labile to a detergent-resistant state

841 late in *T. gondii* daughter cell development (Mann et al., 2002). On the other hand, ubiquitination

of the mother's cytoskeleton marks it for destruction (Silmon de Monerri et al., 2015; Dhara and

843 Sinai, 2016). Although the former informs about maturation and the latter about destabilization,

neither informs directly on the basis of differential stability. In parallel, some of the maternal

845

846 daughters (Ouologuem and Roos, 2014; Periz et al., 2019). An additional mechanism is provided

building blocks (e.g. IMC proteins, glideosome) are recycled in the final growth spurt of the

by differential components present on either mother (e.g. MSC1b (Lorestani et al., 2012),

848 GAP45 (Gaskins et al., 2004), or IMC7, 12, 14, 17, 18 and 20 (Anderson-White et al., 2011;

849 Chen et al., 2015a)) or daughter parasite (e.g. IMC16, 29 (Chen et al., 2015a; Chen et al., 2016)),

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or swapping places from mother to daughter (e.g. SPM3, (Samad et al., 2015)). Such differential
composition of mother and daughters may be factors in differential stability, but to date no single
factor provides a satisfactory explanation.

853 Another potential problem posed by the mother's cytoskeleton during internal budding 854 occurs in the growth phase, where it restricts access to nutrients and complicates the ability to 855 expel waste. This is most relevant for the expanding cell during endopolygeny with large 856 cytoplasmic mass. Although both the apical and basal extremes of the cytoskeleton have 857 openings toward the plasma membrane, they are relatively small and it is not clear whether they 858 are sufficient for the level of exchange needed. However, it is not clear whether the sutures 859 between the alveoli are permeable for diffusion of small molecules, which would void this 860 argument. Either way, a set of apical annuli largely composed of AAP proteins residing in the 861 IMC sutures were recently suggested to function as pores across the IMC (Engelberg et al., 2019; 862 Lentini et al., 2019). In support of this hypothesis is the narrow conservation of AAP protein 863 orthologs only in the Sarcocystidae, indicating their function is most likely in support of internal 864 budding. However, disruption of apical annuli structure resulted in a minor fitness loss of T. 865 gondii tachyzoites, which might be because endodyogeny poses rather limited demands on 866 exchange, especially compared to endopolygenic replication modes where the mother 867 cytoskeleton could pose a much larger barrier. An alternative role for the apical annuli is as a 868 gateway for a burst in dense granule secretion following completion of invasion (Carruthers and 869 Sibley, 1997): electron microscopy studies place dense granule release at this moment in an 870 apical location consistent with the position of the annuli (Dubremetz et al., 1993). Additional 871 studies on the role of the annuli during endopolygeny are needed to firmly differentiate between 872 these two potential roles.

873 In conclusion, we define two principally different asexual cell division modes, external and 874 internal budding, which can both produce as few as two daughters per division round, but in 875 most situations produce many more daughters per division round. The bipartite centrosome 876 model and the identification of several regulators of cell cycle progression and checkpoints in 877 recent years provide a framework to explain these division modes, however the evolutionary 878 history and cell biological features defining, organizing and executing the various division 879 modes are much less clear. Expanding genetic and cell biological toolboxes for parasites 880 representing the various division modes provide exciting future avenues toward resolving the

881	exotic apicomplexan cell division modes and shedding light on the evolutionary pressures that
882	select for diversification and choices for different division modes at different developmental
883	stages.
884	
885	5 Conflict of Interest Statement
886	The authors declare that the research was conducted in the absence of any commercial or
887	financial relationships that could be construed as a potential conflict of interest.
888	
889	6 Author Contributions Statement
890	MG, HW, DH and MD conceived the experiments, HW performed the C. suis experiments, SD
891	and DH performed the S. neurona experiments, MG, CK, AP, BE, and CB performed the
892	Babesia experiments, KE and MG performed SIM microscopy, BE and IC performed electron
893	microscopy, MG wrote the manuscript and all authors edited the manuscript.
894	
895	7 Funding
896	This study was supported by National Science Foundation (NSF) Major Research
897	Instrumentation grant 1626072, National Institute of Health grants AI110690 (MJG), AI110638
898	(MJG), AI128136 (MJG), AI144856 (MJG), and AI128480 (MTD), an American Heart
899	Association pre-doctoral fellowship19PRE34380106 (CDK), a Profillinien start-up grant of the
900	University of Veterinary Medicine Vienna PP16110262 (HLW), an Australian NHMRC CJ
901	Martin fellowship (BE), a post-doctoral fellowship grant 17POST33670577 (KE), a Knights
902	Templar Eye Foundation Career Starter Award (KE), and USDA NIFA grant 2009-65109-05918
903	(DKH). The funders had no role in study design, data collection and analysis, decision to
904	publish, or preparation of the manuscript.
905	
906	8 Acknowledgements
907	We thank Bret Judson and the Boston College Imaging Core as well as Stephan Handschuh and
908	the Imaging Core of the University of Veterinary Medicine Vienna for infrastructure and
909	support, Drs. Naomi Morrissette, Jaime Tarigo, and Jeff Dvorin for discussion, and Drs. David
910	Allred, Kirk Deitsch and Laura Kirkman for sharing reagents.
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912 9 References

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1366 Legend to Figures

1367

1368 Figure 1. Select apicomplexan phylogeny and division modes. 18S ribosomal RNA based 1369 phylogeny of species whose division modes have been studied. Cryptosporidium spp. were used 1370 as outgroup. Bars on the right indicate different naming and biological relationships, with 1371 asexual division modes in blue and red. "in vacuole" and "in cytoplasm" indicate whether 1372 asexual replication occurs in a parasitophorous vacuole, or whether the parasite escapes from the 1373 vacuole and resides in the cytoplasm of the host cell for its replication. Note that only the acute 1374 stage merozoites of S. neurona replicates by endopolygeny in the cytoplasm, whereas metrocytes 1375 preceding the bradyzoites as well as the bradyzoites divide by endodyogeny within a vacuole 1376 supporting a proteoglycan cyst wall. Furthermore, tissue cysts for C. suis have not been described and this lacking ability is likely a secondary loss. In addition, for several Plasmodium 1377 1378 spp. (Simonetti, 1996), as well as *Babesia* and *Theileria* spp. (Jalovecka et al., 2018) it has been shown that sporozoite formation progresses without karyokinesis to produce large polyploid 1379 1380 nuclei while budding is from the cortex. We note that *Plasmodium* sporozoites infect hepatocytes 1381 wherein they divide by schizogony and manipulate the hepatocytes to expand their size, whereas 1382 *Theileria* sporozoites infect white blood cells, replicate by schizogony (Shaw and Tilney, 1992) 1383 and trigger white blood expansion as well as division (i.e. transformation, which basically is 1384 leukemia (Luder et al., 2009; Chakraborty et al., 2017), which contrasts with Babesia sporozoites 1385 as they directly infect red blood cells. "epg with kk" means "endopolygeny with karyokinesis", 1386 in case of *E. callospermophili*; "k.a." means "kinete amplification", in case of select *Babesia* spp. 1387

1388 Figure 2. Apicomplexan asexual cell division by schizogony. A. Schematic representation of 1389 progressive cell division steps during 'classic' schizogony. The mother's cytoskeleton (present in 1390 1) is disassembled following successful host cell invasion resulting in a pleomorphic cell (2) 1391 before onset of mitosis and karyokinesis, which can be asynchronous (3). Daughter cells bud 1392 from the cortex (4/5), positioned by the centrosome anchored at the plasma membrane (4). **B.** 1393 Schematic representation of phases in the cell division during schizogony with limited 1394 karyokinesis as seen during Heamosporidian sprorogenesis. Karyokinesis is inconsistently 1395 following each S/M-phase leading to a nuclei population with varying levels of ploidity. Note 1396 that nuclear cycles within the same nucleoplasm are similar (1), whereas budding is synchronous

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1397 for all nuclei and linked to a final round of S/M (2). Note that the number of offspring in both

1398 forms of schizogony can reach into the 1000s, which is not represented in the schematics.

1399 Chromosome condensation does not occur and the chromosomes are only drawn to convey the

1400 principle of spindle pole attachment. Mother cytoplasm represented in grey, daughter cytoplasms

- 1401 in pink.
- 1402

1403 Figure 3. Binary fission by *Babesia* spp. A-D. *Babesia bigemina* iRBC stages. A.

Immunofluorescence using MAb α-tubulin 12G10 (green) and TgCentrin1(red) polyclonal
antibody. Centrin staining is not observed in interphase (G1) while diffuse and weak during cell

1406 division (cyt.). **B.** Immunofluorescence using MAb α-tubulin 12G10 (green) and microtubule

1407 (+)-end binding protein TgEB1 polyclonal antibody (red) demonstrates that the cortical

1408 microtubules are relatively short and do not reach the nucleus. **C.** Immunofluorescence using

1409 polyclonal guinea pig BbIMC1a antiserum (see Fig S1 for validation) and MAb α-tubulin 12G10

1410 shows the IMC is present during G/M, outlines budding daughters (cyt.), and extends along the

1411 length of the mature G1 parasites. * marks a recently invaded parasites wherein the cytoskeleton

1412 is completing disassembly; we observed many trophozoites without detectable PbIMC1a. **D.**

1413 Transmission electron microscopy of progressive iRBC developmental stages $(1\rightarrow 6)$. Panel 1 is

1414 a disassembling merozoite already escaped from the vacuole displaying typical pattern of 6

1415 numbered remnants of the disassembling mother IMC. Panel 2 represents early daughter

1416 formation ('Mickey Mouse') with the nucleus being separated into the daughter buds. Panels 3,

1417 4, and 5 represent different sections though developing daughters where showing parallel

1418 assembly (in contrast to polar assembly) consistent with the angle under which the daughter buds

1419 assemble. Panel 6 displays two just divided daughters under the typical angle. N, nucleus, SB

spherical body. Yellow marks IMC; orange arrows mark the typical angle and direction of

1421 daughter buds. Scale bars represent 500 nm. E. *Babesia divergens* basal complex upon

1422 completion of cell division. Scale bar represents 500 nm F. Schematic of apicomplexan asexual

1423 cell division by binary fission, which mechanistically represents a binary form of schizogony.

1424 Note that the mother's cytoskeleton is largely disassembled following successful host cell

1425 invasion resulting in a pleomorphic cell (2), but that fragments of the IMC are remaining

1426 throughout the division stages (2-4).

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1427 Figure 4. Apicomplexan asexual cell division by endodyogeny. A-E show progressive steps of

1428 cell division as observed for *T. gondii* tachyzoites. Two daughters bud internally, while the

1429 mother's cytoskeleton is maintained and only destabilized just before emergence of nearly

1430 mature daughters (E). A dark blue SFA fiber anchors the centrosomes in the apex of the daughter

- 1431 buds.
- 1432

1433 Figure 5. Endopolygeny with karyokinesis during merogony of *Cystoisospora suis*. A-D.

1434 Parasites at different stages of endopolygeny. Acetylated (Ac) α-tubulin (red) marks the mother

cells conoid (arrow heads in A, B), subpellicular microtubules, the spindle poles associated with nuclei undergoing mitosis, and the daughter conoids and subpellicular microtubules (C, D).

1437 Antiserum generated against TgNuf2 (green) marks the clustered kinetochores at mitotic

1438 spindles. A. Eight parasites at different stages in the division cycle. Green numbers indicate the

number of spindle poles seen per mother cell, which is expanding non-geometrically in five of

1440 the parasites indicating asynchronous nuclear cycles. **B.** Five parasites at different stages of

1441 endopolygeny are shown. The parasite on the bottom has five nuclei: nuclei marked "M" are in

early stages of mitosis as 2 spindle poles flanking a single 2N kinetochore cluster are seen; the

1443 circled nuclei represent just completed mitosis as the kinetochore clusters are relatively weak

1444 intensity (1N vs. 2N) and associated with only a single spindle pole. **C**, **D**. Two examples of cells

1445 at the synchronous daughter budding stage, which both produce 12 daughters as enumbered by

1446 the number of kinetochore clusters, again, consistent with non-geometric expansion of the nuclei.

1447 E. Schematic of apicomplexan asexual cell division by endopolygeny with karyokinesis. Note

1448 that the mitotic cycles of the nuclei in the same cytoplasm are not synchronous (2) resulting in

non-geometrically expanded daughter numbers (4-5). Also note that the mother's cytoskeleton is

1450 maintained throughout the meront (technically 'schizont') stages and is only destabilized just

- 1451 before emergence of nearly mature daughters (5).
- 1452

1453 Figure 6. Endopolygeny without nuclear fission by *Sarcocystis neurona* in the intermediate

host. A. Staining of progressive cell division cycle stages (1-4) with α-tubulin (green) marking

1455 the subpellicular microtubules of the mother cytoskeleton (arrow heads), the mitotic spindles (2b,

right parasite; 3) and daughter merozoites subpellicular microtubules (4). Centrin staining (red)

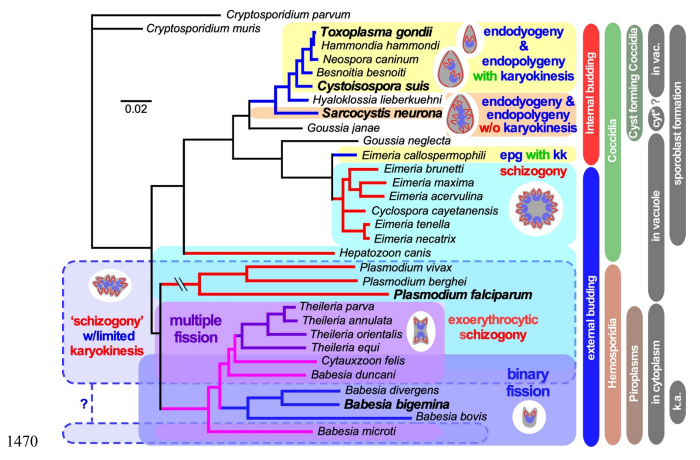
1457 marks the centrosomes, which due to z-stack selection are not visible for all spindles/parasites.

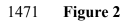
Comparative apicomplexan cell division

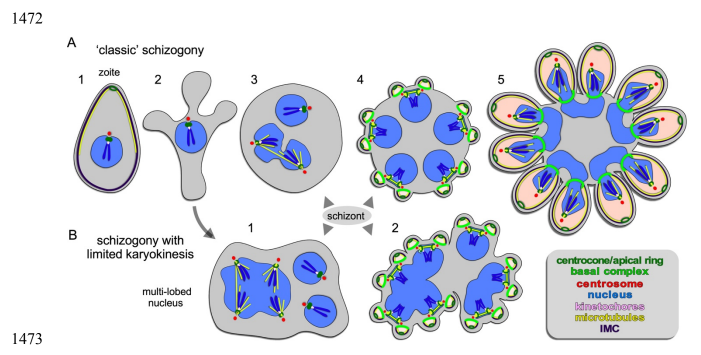
- 1458 Scale bar applies across panels. **B.** Overexpression of a *Toxoplasma* YFP-IMC15 fusion protein
- 1459 (green) highlights the mother cell's cortical IMC sutures in both panels, whereas in further
- 1460 progressed panel 2 the bright internal spots mark the centrosomes poised for budding.
- 1461 Arrowhead marks the apical end of the mother parasite. Panel B modified from (Dubey et al.,
- 1462 2017). Scale bar applies across panels. **C.** Schematic of apicomplexan asexual cell division by
- 1463 endopolygeny without karyokinesis. Note the polyploid nuclei undergoing synchronized cycles
- 1464 of M-phase and mitosis resulting in geometrically expanded daughter cell numbers. Also note
- 1465 that the mother's cytoskeleton is maintained throughout the schizont stages (2-4) and is only
- 1466 destabilized just before emergence of nearly mature daughters (5).

Comparative apicomplexan cell division





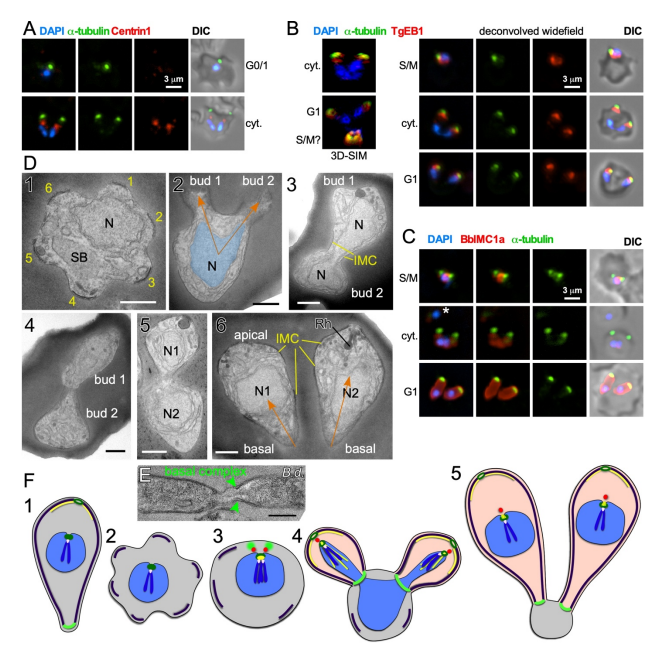


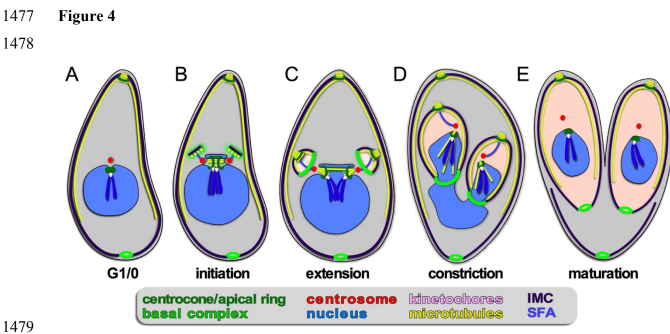


Comparative apicomplexan cell division

1474 Figure 3

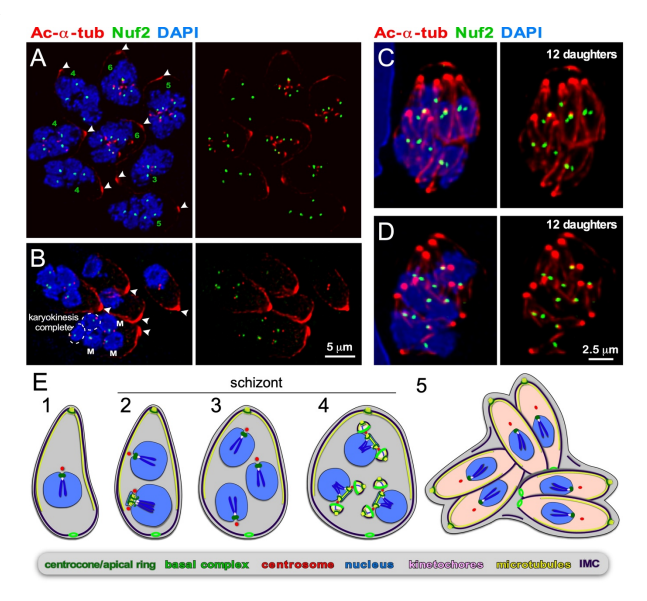
1475





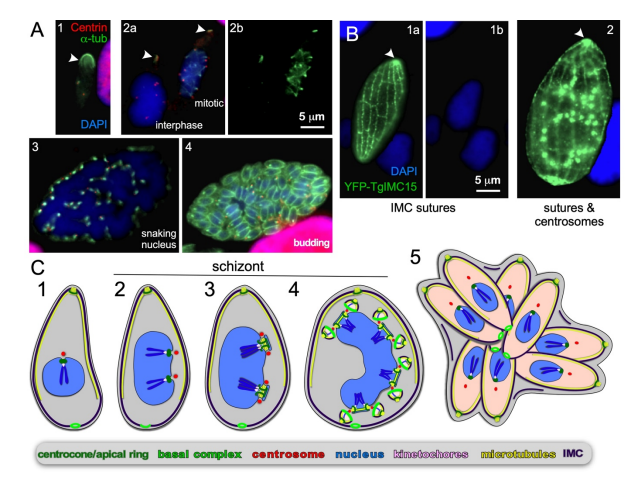
Comparative apicomplexan cell division

Figure 5



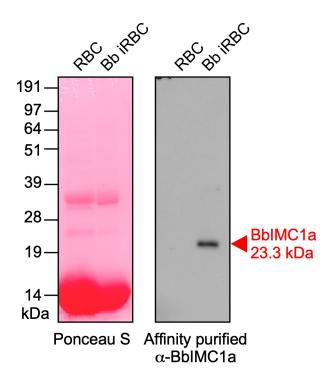
Comparative apicomplexan cell division

Figure 6

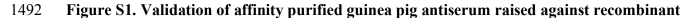


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- 1486 Supplementary Material
- 1487
- 1488 Figure S1
- 1489



1490 1491



1493 His6-BbIMC1a by western blot. Left panel: PonceauS staining of the western blot serving as

1494 loading control; Right panel: serum affinity purified against recombinant His6-BbIMC1a diluted

- 1495 1:250. RBC indicates cow red blood cell total lysate; Bb iRBC indicates total lysate of cow red
- 1496 blood cells with a *B. bigemina* parasitemia of 12%. Equal amounts of lysate were loaded across
- 1497 lanes. The predicted MW of BbIMC1a is 23.3 kDa.